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Post Authorisation Assessments

Frontline Plus Spot-on Dog S Vm 08327/4270

13 May 2025	Deletion of a non-significant specification parameter for an excipient. Deletion of a non-significant specification parameter for an active
	substance.
18 February 2025	Change in the batch size of the finished product.
18 February 2025	Minor change in the manufacturing process of the finished product.
18 January 2025	Change in the specification parameters and/or limits of the finished product: - Other changes Change in the specification parameters and/or limits of the finished product: - Change outside the approved specifications limits range
13 April 2023	Change in the name or address or contact details of a qualified person for pharmacovigilance.
16 June 2021	Addition to a test procedure for the finished product.
08 March 2021	Change in the number of units (pipettes) in a pack within the range of the currently approved pack sizes of the finished product.
	Change to part of the (primary) packaging material not in contact with the finished product formulation.
15 October 2020	Addition of a site where batch control/testing takes place.
19 August 2020	Renewal – national.
28 May 2020	Change in the name of a manufacturer of the finished product, also responsible for batch release.
12 March 2020	Change in legal distribution category from NFA-VPS to AVM-GSL.
05 November 2019	Change in the safety database of an existing pharmacovigilance system as described in the DDPS.
08 October 2019	Changes in the qualitative and quantitative composition of the immediate packaging of the finished product.
15 April 2019	Update to the ASMF
12 February 2019	Change in the name and address of the manufacturer of the finished product.
16 November 2018	Change in the name and address of the marketing authorisation holder from Merial Animal Health Limited, Sandringham House, Sandringham Avenue, Harlow Business Park, Harlow, Essex, CM19 5TG, United Kingdom to Boehringer Ingelheim Animal Health UK Limited, Ellesfield Avenue, Bracknell, Berkshire, RG12 8YS, United Kingdom.
04 October 2018	Change in the address of a manufacturer of the active substance where no Ph. Eur. Certificate of Suitability is part of the approved

	dossier.
	Deletion of a manufacturing site for an active substance.
29 August 2018	Change in the address of the supplier used for the manufacture
	of the active substance.
	Addition of a new specification parameter with its corresponding
	test method of an active substance used in the manufacturing
	process of the active substance.
	Change of specification(s) of a former non Pharmacopoeial
	active substance to comply with the Ph. Eur. or with a national
	pharmacopoeia of a Member State.
26 February 2018	Minor changes to an approved test procedure of the finished
	product.
	Minor change in the manufacturing process of the finished
	product.
27 November 2017	Changes to the labelling and/or package leaflet.
28 April 2017	Extension of a re-test period of the active substance.
10 February 2017	Approval of mock ups
10 March 2016	To change the legal category from POM-V to NFA-VPS
14 January 2016	Change in the QPPV and/or QPPV contact details and/or back-
	up procedure
22 December 2015	Change of invented product name